Hybridization was introduced to explain molecular structure when the valence bond theory failed to correctly predict them. It is experimentally observed that bond angles in organic compounds are close to 109°, 120°, or 180°. According to Valence Shell Electron Pair Repulsion (VSEPR) theory, electron pairs repel each other and the bonds and lone pairs around a central atom are generally separated by the largest possible angles.

**Introduction**

Carbon is a perfect example showing the need for hybrid orbitals. As you know, Carbon's ground state configuration is:

![Carbon Electron Configuration](image)

According to Valence Bond Theory, carbon should form two covalent bonds, resulting in a CH₂, because it has two unpaired electrons in its electronic configuration. However, experiments have shown that CH₂ is highly reactive and cannot exist outside of a reaction. Therefore, this does not explain how CH₄ can exist. To form four bonds the configuration of carbon must have four unpaired electrons.

The only way CH₄ it can be explained is is, the 2s and the 3 2p orbitals fused together to make four, equal energy sp³ hybrid orbitals. That would give us the following configuration:

![Carbon Hybrid Configuration](image)

Now that carbon has four unpaired electrons it can have four equal energy bonds. The hybridization of orbitals is also greatly favored because hybridized orbitals are lower in energy compared to their separated, unhybridized counterparts. This results in more stable compounds when hybridization occurs. Also, major parts of the hybridized orbitals, or the frontal lobes, overlap better than the lobes of unhybridized orbitals. This leads to better bonding.

The next section will explain the various types of hybridization and how each type helps explain the structure of certain molecules.

**sp³ hybridization**

sp³ hybridization can explain the tetrahedral structure of molecules. In it, the 2s orbitals and all three of the 2p orbitals hybridize to form four sp orbitals, each consisting of 75% p character and 25% s character. The frontal lobes align
themselves in the manner shown below. In this structure, electron repulsion is minimized.

**Energy changes occurring in hybridization**

\[
\begin{array}{c}
\text{2s} \\
\text{1s}
\end{array} \rightarrow \begin{array}{c}
\text{1s} \\
\text{2p}
\end{array} \rightarrow \text{sp}^3 \text{ hybrid}
\]

**Edit section**

Hybridization of an s orbital with all three p orbitals \((p_x, p_y, \text{and } p_z)\) results in four \(\text{sp}^3\) hybrid orbitals. \(\text{sp}^3\) hybrid orbitals are oriented at bond angle of 109.5° from each other. This 109.5° arrangement gives tetrahedral geometry (Figure 4).

**Example: \(\text{sp}^3\) Hybridization in Methane**

Because carbon plays such a significant role in organic chemistry, we will be using it as an example here. Carbon's 2s and all three of its 3p orbitals hybridize to form four \(\text{sp}^3\) orbitals. These orbitals then bond with four hydrogen atoms through \(\text{sp}^3\)-s orbital overlap, creating methane. The resulting shape is tetrahedral, since that minimizes electron repulsion.

**Hybridization**

\[
\begin{array}{c}
\text{C} \\
\text{H}_2\text{O}
\end{array}
\]

**Lone Pairs:** Remember to take into account lone pairs of electrons. These lone pairs cannot double bond so they are placed in their own hybrid orbital. This is why \(\text{H}_2\text{O}\) is tetrahedral. We can also build \(\text{sp}^3d\) and \(\text{sp}^3d_2\) hybrid orbitals if we go beyond s and p subshells.
**sp\(^2\) hybridization**

sp\(^2\) hybridization can explain the trigonal planar structure of molecules. In it, the 2s orbitals and two of the 2p orbitals hybridize to form three sp orbitals, each consisting of 67% p and 33% s character. The frontal lobes align themselves in the trigonal planar structure, pointing to the corners of a triangle in order to minimize electron repulsion and to improve overlap. The remaining p orbital remains unchanged and is perpendicular to the plane of the three sp\(^2\) orbitals.

**Energy changes occurring in hybridization**

Hybridization of an s orbital with two p orbitals (p\(_x\) and p\(_y\)) results in three sp\(^2\) hybrid orbitals that are oriented at 120° angle to each other (Figure 3). Sp\(^2\) hybridization results in trigonal geometry.

**Example: sp\(^2\) Hybridization in Aluminum Trihydride**

In aluminum trihydride, one 2s orbital and two 2p orbitals hybridize to form three sp\(^2\) orbitals that align themselves...
in the trigonal planar structure. The three Al sp² orbitals bond with 1s orbitals from the three hydrogens through sp²-s orbital overlap.

Example: sp² Hybridization in Ethene

Similar hybridization occurs in each carbon of ethene. For each carbon, one 2s orbital and two 2p orbitals hybridize to form three sp² orbitals. These hybridized orbitals align themselves in the trigonal planar structure. For each carbon, two of these sp orbitals bond with two 1s hydrogen orbitals through s-sp orbital overlap. The remaining sp² orbitals on each carbon are bonded with each other, forming a bond between each carbon through sp²-sp² orbital overlap. This leaves us with the two p orbitals on each carbon that have a single carbon in them. These orbitals form σ bonds through p-p orbital overlap, creating a double bond between the two carbons. Because a double bond was created, the overall structure of the ethene compound is linear. However, the structure of each molecule in ethene, the two carbons, is still trigonal planar.
**sp Hybridization**

sp Hybridization can explain the linear structure in molecules. In it, the 2s orbital and one of the 2p orbitals hybridize to form two sp orbitals, each consisting of 50% s and 50% p character. The front lobes face away from each other and form a straight line leaving a 180° angle between the two orbitals. This formation minimizes electron repulsion. Because only one p orbital was used, we are left with two unaltered 2p orbitals that the atom can use. These p orbitals are at right angles to one another and to the line formed by the two sp orbitals.

**Energy changes occurring in hybridization**

![Figure 1: Notice how the energy of the electrons lowers when hybridized.](image)

These p orbitals come into play in compounds such as ethyne where they form two addition bonds, resulting in a triple bond. This only happens when two atoms, such as two carbons, both have two p orbitals that each contain an electron. An sp hybrid orbital results when an s orbital is combined with p orbital (Figure 2). We will get two sp hybrid orbitals since we started with two orbitals (s and p). sp hybridization results in a pair of directional sp hybrid orbitals pointed in opposite directions. These hybridized orbitals result in higher electron density in the bonding region for a sigma bond toward the left of the atom and for another sigma bond toward the right. In addition, sp hybridization provides linear geometry with a bond angle of 180°.

**Example: sp Hybridization in Magnesium Hydride**

In magnesium hydride, the 3s orbital and one of the 3p orbitals from magnesium hybridize to form two sp orbitals. The two frontal lobes of the sp orbitals face away from each other forming a straight line leading to a linear structure. These two sp orbitals bond with the two 1s orbitals of the two hydrogen atoms through sp-s orbital overlap.

**Example: sp Hybridization in Ethyne**

The hybridization in ethyne is similar to the hybridization in magnesium hydride. For each carbon, the 2s orbital hybridizes with one of the 2p orbitals to form two sp hybridized orbitals. The frontal lobes of these orbitals face away...
from each other forming a straight line. The first bond consists of sp-sp orbital overlap between the two carbons. Another two bonds consist of s-sp orbital overlap between the sp hybridized orbitals of the carbons and the 1s orbitals of the hydrogens. This leaves us with two p orbitals on each carbon that have a single carbon in them. This allows for the formation of two π bonds through p-p orbital overlap. The linear shape, or 180° angle, is formed because electron repulsion is minimized the greatest in this position.

Hybridization

References
2. Francis A. Carey *Advanced Organic Chemistry* Springer 2001. 4-6
Problems

Using the [Lewis Structures](#), try to figure out the hybridization (sp, sp\(^2\), sp\(^3\)) of the indicated atom and indicate the atom's shape.

1. The carbon.

   ![Lewis Structure of Carbon](#)

   \[ \text{H-C-H} \]

   The carbon has no lone pairs and is bonded to three hydrogens so we just need three hybrid orbitals, aka sp\(^2\).

2. The oxygen.

   ![Lewis Structure of Oxygen](#)

   \[ \text{O-H} \]

   Don't forget to take into account all the lone pairs. Every lone pair needs its own hybrid orbital. That makes three hybrid orbitals for lone pairs and the oxygen is bonded to one hydrogen which requires another sp\(^3\) orbital. That makes 4 orbitals, aka sp\(^3\).

3. The carbon on the right.

   ![Lewis Structure of Carbon](#)

   \[ \text{H-C-C-H} \]

   The carbon is bonded to two other atoms, that means it needs two hybrid orbitals, aka sp.

Answers

1. sp\(^2\) - Trigonal Planar

   The carbon has no lone pairs and is bonded to three hydrogens so we just need three hybrid orbitals, aka sp\(^2\).

2. sp\(^3\) - Tetrahedral

   Don't forget to take into account all the lone pairs. Every lone pair needs its own hybrid orbital. That makes three hybrid orbitals for lone pairs and the oxygen is bonded to one hydrogen which requires another sp\(^3\) orbital. That makes 4 orbitals, aka sp\(^3\).

3. sp - Linear

   The carbon is bonded to two other atoms, that means it needs two hybrid orbitals, aka sp.
An easy way to figure out what hybridization an atom has is to just count the number of atoms bonded to it and the number of lone pairs. Double and triple bonds still count as being only bonded to one atom. Use this method to go over the above problems again and make sure you understand it. It's a lot easier to figure out the hybridization this way.

Contributors

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